PATENT SPECIFICATION

1 549 180 (11)

(21) Application No. 29854/75

(22) Filed 16 July 1975 (22) Filed 4 June 1976

(21) Application No. 23171/76 (21) Application No. 23174/76

(22) Filed 4 June 1976

(23) Complete Specification filed 16 July 1976

(44) Complete Specification published 25 July 1979

(51) INT CL2 C08L 83/04 C08K 5/17 D06 M 13/46 15/66

(52) Index at acceptance

5

10

15

20

25

30

35

C3T 6D2A 6D4B 6F2 6H1 6H4X D1P 21Y 23Y 240 24Y 272 27Y 36Y 370 52Y 53Y 541 543 54Y 580 581 584 587 595 59Y 610 621 62Y 636 65Y 660 66Y 670 70Y 744 74Y A18 B2B1 B5B C2A10 C2A12A10 C2A12A9 C2C16 C2C2 C2C3 C2C5 C2C7 C2C8 C2C9 C2CX L2 L6

(72) Inventors RICHARD JOHN DUMBRELL

JEAN PIERRE CHARLES IGNACE MARIE LECLERCO, RITA MARIE ANTOINETTE DE BAKKER PIERRE CHARLES EMILE GOFFINET **BRIAN ANTHONY BROWN** RONALD EDWARD ATKINSON FREDERICK EDWARD HARDY

(54) TEXTILE TREATING COMPOSITIONS

We, THE PROCTER & GAMBLE COMPANY, a Company organised under the laws of the State of Ohio, United States of America, of 301, East Sixth Street, Cincinnati, Ohio 45202, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following

The invention relates to compositions and a method for treating fabrics in an aqueous bath such as the final rinse after a washing process to improve various properties of the fabric.

It has been known for several years that fabrics can be made to feel softer by treatment in a dilute solution or dispersion of certain cationic quaternary ammonium derivatives, and rinse-additive compositions have been marketed for this purpose.

The present invention surprisingly makes it possible to provide substantial additional benefits by an unexpected combination of fabric conditioning agents. These benefits may include some or all of: easier ironing, anti-static properties, pleasanter feel of the fabrics, soil release properties. It appears that recognition of improved "ease of ironing" can arise from a combination of at least three factors, namely fewer wrinkles to be removed, wrinkles more easily removed (e.g. with less weight upon the iron), or more completely removed, and less effort required to slide the iron along the fabric. "Pleasanter" feel can be consistently observed by experienced judges, although it is not easy to define in words the sensation or combination of sensations they like. Anti-static and soil release properties improve the achieving and maintaining of soil-free fabrics.

The combination of agents of the invention involves the presence of both a cationic compound having one or more long chain alkyl groups and a silicone, as further defined hereinbelow.

Silicones have been used to provide various benefits, for instance as waterproofing agents and as "ironing aids", but have usually been applied to the fabrics, during manufacture or during make-up of articles of clothing, in the form of relatively concentrated dispersions or solutions of silicones, either by processes such as padding, or by spray on. In other words, the prior art treatments have relied upon treatment with relatively concentrated, e.g. 2-3% by weight and upwards, dispersions or solutions of silicones to ensure that some silicone remained upon the surface of the fabrics. Often, especially for water-proofing treatment, the fabric was subsequently treated with catalysts to cause cross linking or setting of the silicone. If normal commercial silicones are applied to fabrics from dilute aqueous systems, they are not substantive





5

10

15

20

25

30

35

2

5

10

15

20

25

30

35

40

45

5

10

15

20

30

35

40

45

to a useful degree, that is insufficient silicone is present in the dilute residual liquor in the fabric to provide any appreciable effect, and therefore the advantageous effects of the silicone on the fabrics are greatly reduced or lost.

It has now surprisingly been found that if a silicone is incorporated, even at very low concentrations, in a bath containing certain fabric substantive cationic textile softeners, the silicone and softener being used in certain proportions, the silicone, together with the textile softener, tends to migrate to the surface of the fabric and becomes substantive, and concentrated thereon. Thus, there can be achieved a very desirable overall effect involving both softening benefits and such additional benefits as mentioned hereinabove. These latter can be varied by varying the silicone.

According to the present invention there is provided a textile treating composition in the form of an aqueous dispersion comprising:

a fabric substantive cationic compound selected from

quaternary mono-ammonium compounds having either two C12-C20 alkyl chains or one C₁₈—C₂₄ alkyl chain;

quaternary imidazolinium textile softeners;

polyammonium compounds of the general formula

wherein R_8 is a straight or branched chain C_{10} — C_{22} alkyl group or alkenyl group; R_{10} , R_{11} and R_{12} are independently selected from hydrogen, C_1 — C_3 alkyl, $(C_2H_4O)_0H$ or $(C_3H_6O)_0H$, wherein p+q is $\ll 25$; R_0 is defined as R_0 or as R_{11} and R_{12} ; m is from 0 to 8, n is from 2 to 6 and X^{\to} is an anion;

(iv) compounds of the general formula:

$$R_{13} - N + - \left[\begin{pmatrix} R_{13} \\ + \\ R_{13} \end{pmatrix} + \begin{pmatrix} R_{13} \\ + \\ R_{13} \end{pmatrix} + \begin{pmatrix} R_{13} \\ + \\ R_{13} \end{pmatrix} \right] R_{13}, \quad (s+1) \times \Theta$$
25

wherein R_{13} is hydrogen or a C_1 — C_4 alkyl group, r is an integer from 2 to 6, s is not less than 3 and X^{\ominus} is an anion; and

mixtures thereof; and

an emulsion of a predominantly linear di(C1-C5) alkyl or alkyl-aryl siloxane in which the alkyl groups may be partially or wholly fluorinated and which may be substituted with cationic nitrogen groups, the siloxane having a viscosity at 25°C of at least 100 centistokes and up to 8000 centistokes; the weight ratio of the siloxane content of component (b) to component (a) being in the range from 5:1 to 1:100.

The weight ratio of the siloxane content of component (a) preferably lies in the range from 2:1 to 1:10, especially from 1:1 to 1:3,

Fabric substantive cationic component

The cationic compounds suitable include those commonly used in rinse-added textile softening compositions. They include quaternary ammonium salts of general formula

wherein either (a) R2 and R3 (which may be the same or different) represent methyl, ethyl, propyl or benzyl, and either R and R1 each represent a straight or branched chain alkyl group having 12 to 20 carbon atoms, or R represents a straight or branched chain alkyl group having 18 to 24 carbon atoms and R₁ represents methyl, ethyl, propyl or benzyl; or (b) R₂ and R₃ together with the nitrogen atom form a 5membered or 6-membered heterocyclic ring and R represents a straight or branched

45

3	1,549,180	3
	chain alkyl group having 18 to 24 carbon atoms; and X- is an anion. The long chain alkyl groups may be derived from natural fats e.g. coconut, or more preferably tallow,	
	or from petroleum or synthetically. In a preferred group of salts of formula (I), R and R ₁ each represent an alkyl	_
5	group having 16 to 18 carbon atoms, R2 and R3 each represent methyl, and X-represents Cl-, Br or OSO ₃ CH ₃	5
	Other anions include nitrite, acetate and phosphate. Specific examples of particularly preferred cationic softening agents include the	
10	following: —-tallowtrimethyl ammonium chloride,	10
	—tallowdimethyl (3-tallowalkozypropyl) ammonium chloride, —ditallow dimethyl ammonium chloride,	
	-ditallow dimethyl ammonium methyl sulphate,	
15	-eiicosyltrimethyl ammonium chloride, and -dieicosyldimethyl ammonium chloride.	15
15	Examples of other suitable cationic softening agents suitable for use in the	
	invention herein include the following: —ditetradecyldimethyl ammonium chloride,	
20	—dipentadecyldimethyl ammonium chloride, —didodecyldiethyl ammonium chloride,	20
20	didodecyldipropyl ammonium chloride,	
	—ditetradecyldiethyl ammonium chloride, —ditetradecyldipropyl ammonium chloride,	
25	—ditallowdiethyl ammonium chloride, —ditallow dipropyl ammonium chloride,	25
23	—tallowdimethyl benzyl ammonium chloride,	
	—tallowdiethyl benzyl ammonium chloride, —didodecyldiethyl ammonium acetate,	
30	—tallowtrimethyl ammonium acetate, —tallowdimethyl benzyl ammonium nitrite, and	30
30	dirallowdipropyl ammonium phosphate.	
	Other cationic softening agents of formula (1) are known and include variations wherein R and R ₁ can also represent a phenyl radical or a hydroxy-substituted alkyl	
35	of 1, 2 or 3 carbon atoms. Many other cationic quaternary ammonium softening agents, which are useful	35
33	berein are known: for example, alkyl [C ₁₂ to C ₂₀]-pyridinium chlorides, alkyl	
	[C ₁₂ to C ₂₀]-alkyl [C ₁ to C ₃]-morpholinium chlorides, and quaternary derivatives of amino acids and amino esters.	
40	Cationic quaternary imidazolinium compounds are also suitable as softening agents in the compositions herein. The structure of these compounds is not fully	. 40
40	established but is believed to be either	
	- J	

wherein R₄ represents hydrogen or alkyl having 8 to 25, preferably at least 15, carbon atoms, R₅ represents alkyl having 1 to 4, preferably 1 or 2, carbon atoms, R₆ represents alkyl having 1 to 4 carbon atoms or hydrogen, R₇ represents alkyl having 8 to 25, preferably at least 15, carbon atoms and X⁻ is an anion, preferably methyl sulphate or chloride. Other suitable anions include bromide, acetate, nitrite and phosphate.

Particularly preferred are those compounds of formula (II) in which both R, and R, represent alkyl having 16 to 25 (especially 16 to 18 or 20 to 22) carbon atoms.

A further group of fabric substantive cationic compounds suitable for use in the present invention is characterised by the following general formula

$$R_{B} - \frac{R_{1}^{Q}}{R_{1}} - co_{12}I_{1} - \frac{R_{1}^{Q}}{R_{12}} - \left[co_{12}I_{1} - \frac{R_{1}^{Q}}{R_{12}}\right]_{m} \left[x^{-}\right]$$
(III) 5

wherein R_8 is a straight or branched chain C_{10} — C_{22} alkyl group or alkenyl group, R_{10} , R_{11} and R_{12} are independently selected from hydrogen, C_1 — C_3 alkyl, $(C_2H_4O)_0H$ and $(C_3H_6O)_0H$ wherein p+q is $\ll 25$ and wherein not all of the R_{10} , R_{11} and R_{12} groups are C_1 — C_3 alkyl, R_9 is defined as R_8 or as R_{10} , R_{11} and R_{12} , m is from 0 to 8, n is from 2 to 6 and $X^{(-)}$ represents one or more anions having a total charge balancing that of the nitrogen atoms. (Highly preferred compounds of this group are the acid salts of diamine compounds where m=0 in formula (III) above). Particularly preferred salts of diamine compounds where m=o in formula (III) above). Particularly preferred compounds within this group are N-C₁₆₋₁₈ alkyl N,N,N¹,N¹,N¹,N¹, pentamethyl-1,3-propane diamine diacetate and dichloride (i.e. where $R_8 = C_{16} - C_{18}$ alkyl R_9 R_{10} R_{11} R₁₂=CH₃ m=0 X=Cl or CH₃COO— in III above).

Another preferred class of compounds within the above group are the fabric softening polyamine salts, i.e. those compounds of formula (III) above in which at least N,N'N' triethanol-1,3-propane diamine dichloride or diacetate (i.e. where R₆=C₁₆—C₁₈ alkyl R₀ R₁₁-C₂H₄OH R₁₀ and R₁₂=hydrogen m=0 and x=Cl⁻ or CH₃COO in III

above) is an example of this class. Acid salts of diamine compounds, as used herein, are the addition products

formed when certain acids add to the amino moieties of the diamines and form monoor di-ammonium salts.

The diamine acid salts can be partially acidified diamine salts (i.e. only one nitrogen atom quaternized with acid) of fully acidified diamine salts (i.e. both nitrogen atoms quaternized with acid).

A variety of acids can be used herein to form the acid salts so long as the anion of the diamine acid salt formed is stable under fabric rinse conditions and is noninterfering in the rinse. Suitable acids include organic and inorganic acids such as hydrochloric acid, =acetic acid, sulphuric acid, lactic acid, stearic acid, formic acid, citric acid and a large variety of others. Particularly preferred acids used to form diamine acid salts include acetic acid and hydrochloric acid.

Nonlimiting examples of such diamine acid salts include:

35 $[C_{16}H_{33}-NH(CH_3)-(CH_2)_3-NH(CH_3)_2]^{++}(CH_3COO)_2^{--}$ 35

 $[C_{15}H_{37}-NH(CH_3)-(CH_2)_2-NH(C_2H_5)_2]^{++}(CH_5COO)_2^{-}$

 $[C_{12}H_{25}-NH(CH_3)-(CH_2)_3-H_2N-C_{12}H_{25}]^{++}Cl_2^{-}$

 $[C_{12}H_{23}-NH(C_2H_3)-(CH_2)_3-NH(C_3H_7)_2]^{++}(CH_3SO_4)_2^{--}$

 $[R_{Tallow} NH_2 - (CH_2)_3 - NH(C_2H_5)_2]^{++} Br_2$

10

15

20

25

30

 $[C_{20}H_{43}-NH(CH_3)-(CH_2)_2-NH(CH_3)_2]^{4+}Cl_2$ 40 40

 $[C_{13}H_{31}-NH(C_{2}H_{5})-(CH_{2})_{3}-NH_{3}]^{++}(CH_{3}COO)_{2}$

 $[C_{1s}H_{sr}-NH_{z}-(CH_{z})_{s}-H_{z}N-CH_{s}]$ (HCOO),

 $[C_{16}H_{23}-NH_2-(CH_2)_3-H_2N-C_{16}H_{33}]^{++}Cl_2^{-}$

 $[R_{Tallow} NH(CH_3)-(CH_2)_3-NH(C_2H_3)_2]^{++}(CH_3COO)_2^{-}$

45 $[C_{10}H_{33}NH(CH_3)-(CH_2)_5-NH(C_2H_3)_5]^+(CH_3SO_4)_5$

 $[C_{12}H_{25}NH(C_2H_5)-(CH_2)_2-NH(C_3H_7)_2]^{++}$ Cl_2 - and

 $[C_{14}H_{29}NH(CH_3)-(CH_2)_5-(CH_1)NH-(C_8H_{17})]$ + $(CH_5COO)_2$ wherein in the above formulae R_{Tallow} is the alkyl group derived from tallow fatty acid. 10

15

20

25

30

45

5

10

15

20

25

30

35

40

45

50

55

Other examples of suitable compounds include those in which the starting diamine is N-tetradecyl, N'-propyl-1,3-propane-diamine; N-eicosyl,N,N',N'-triethyl-1,2-ethane-diamine and N-octadecyl, N,N',N'-tripropyl-1,3-propane-diamine.

The form in which the diamine acid salt is derived is not critical. The diamine acid salt can be formed from diamines in situ during the preparation of the aqueous fabric conditioning compositions herein or can alternatively, for example, be obtained

in the compositions of the present invention.

The fully acidified diamine salts (i.e. the diacid salts) are particularly preferred inasmuch as these materials permit formation of highly stable, clear, aqueous liquid fabric conditioning compositions.

as the acid salt from commercial sources (e.g. Duomac T marketed by Armour-Hess Co). Mixtures of diamine acid salts can, of course, be used as the static control agent

Preferred aqueous fabric conditioning compositions contain diamine acid salts wherein R₁ is alkyl of from about 16 to about 18 carbon atoms; and R₂, R₃ and R₄ are hydrogen or alkyl of from about 1 to about 2 carbon atoms and n is 2 or 3, more preferably 3.

Diamine acid salts useful in the invention herein are commercially available under a variety of trade marks including Duomeens and Duomacs (marketed by Armour-Hess Co) and Dinozem and Dinoremac (marketed by CECA/Pierrefitte-Auby). Moreover, the starting diamine compounds can be prepared in accordance with processes disclosed in the art, as, for example, in U.S. Patent 2,267,205 issued December 23, 1941 to Kyrides and U.S. Patent 2,246,524 issued June 24, 1941 to Kyrides.

Also within the scope of the polymeric compounds embraced by formula (III) are fully substituted, i.e. quaternised compounds. As indicated above, these polyquaternary compounds do not consist wholly of C₁—C₃ alkyl substituent groups but also contain a level of polyalkenoxy substitution which may be polyethenoxy, polypropenoxy or mixtures of the two and in which each poly alkenoxy chain consists of not more than 25 units.

Polyalkylene imine salts have also been found to be of value in the present invention. These salts have the general formula:

wherein R_{1s} is hydrogen or a C₁—C_s alkyl group, r is an integer from 2—6, preferably from 2—4 and most preferably 2, s is not less than 3, preferably from 6—24 and X is as defined above. Preferably R_{1s} is methyl and s has a value from 8 to 16.

Silicone Component

10

15

20

25

30

35

40

45

50

55

As mentioned previously, the silicone component is an aqueous emulsion of a predominantly linear polydialkyl or alkyl, aryl siloxane in which the alkyl groups can have from one to five carbon atoms and may be wholly or partially fluorinated. Suitable silicones are polydimethyl siloxanes having a viscosity at 25°C in the range from 100 to 8,000 centistokes preferably in the range from 1000 to 5000 centistokes.

As recited hereinbefore, the combination of a fabric substantive quaternary ammonium textile softening compound and a silicone of the above type materially improves the substantivity of the latter. It is postulated that this enhanced substantivity arises from a 'carrier' effect by means of which the positivity charged fabric softener molecules associate with the silicone molecules and cause them to migrate to the fabric surface. However, experiments have shown that the distribution of this combination on the fabric is less than optimum i.e. some areas of fabric receive a high concentration of silicone while others receive very little if any silicone.

It has however been found that the ionic charge characteristics of the silicone as used in the combination are important in determining both the extent of deposition and the evenness of distribution of the silicone and hence the properties of a fabric treated therewith.

Silicones having cationic character show an enhanced tendency to deposit.

As mentioned above, silicones found to be of value in providing fabric feel benefits have a predominantly linear character and are preferably polydialkyl siloxanes in which the alkyl group is most commonly methyl. Such silicone polymers are frequently manufactured commercially by emulsion polymerisation using a strong acid or strong alkali catalyst in the presence of a nonionic or mixed nonionic-anionic emulsifier system.

6	29.75,200	
	In combination with the cationic fabric softening agent, anionic- or nonionic emulsified silicone polymers tend to aggregate in dilute aqueous solution due to the attraction between the negatively or non-charged emulsifier and the positively charged fabric softening agent. However, the provision of a silicone emulsion having a like	e
5	charge to that of the fabric softening agent would tend to reduce this effect and a further reduction might be expected from the tendency of the charged silicone-droplets to repel each other. Thus, in the present invention, the silicone component embraces a silicone of cationic character which is defined as being one of	5
10	 (a) a predominantly linear di C₁—C₅ alkyl or C₁—C₅ alkyl, aryl siloxane, prepared by emulsion polymerisation using a cationic surfactant as emulsifier. (b) an α₃ω-di quaternised di C₁—C₅ alkyl or C₁—C₅ alkyl, aryl siloxane 	10
15	(c) an amino-functional di C ₁ —C ₅ alkyl or alkyl aryl siloxane polymer in which the amino group may be substituted and may be quaternised and in which the degree of substitution (d.s.) lies in the range 0.001 to 0.1, preferably .01—0.075. provided that the viscosity at 25°C of the silicone is from 100 to 8,000 cs.	15
	a) Cationic emulsion polymerised siloxanes	
20	Cationic emulsion polymerised siloxanes are known in the art and can be prepared by strong alkali or acid catalysis of siloxane monomer(s) in the presence of a cationic emulsifying agent. Hyde and Wehryl U.S. Patent No. 2,891,920 describes general procedures for such polymerisations and Examples 1—6 of the patent provide specific teaching of the required reaction conditions. The siloxane monomer can be	20
25	any di lower alkyl siloxane such as dimethyl, diethyl dipropyl, or ethyl butyl siloxane or alkyl, aryl siloxane such as methyl, phenyl siloxane or ethyl phenyl siloxane. However, the preferred starting material for emulsion polymerisation is normally a cyclic trimer or tetramer of the desired siloxane. The emulsifying agent can be any one of a wide range of cationic surfactants	25
30	ench as:	30
	Aliphatic fatty amines and their derivatives such as dodecylamine acetate, octa- decylamine acetate and acetates of the amines of tallow fatty acids; homologues of aromatic amines having fatty chains such as dodecylaniline; heterocylic aliphatic digratives such as undecylimidazoline; fatty amines derived from disub-	25
35	stituted amines such as oleylaminodiethylamine; derivatives of ethylene diamine; quaternary ammonium compounds such as dioctadecyldimethyl ammonium chloride, didodecyldimethyl ammonium chloride and dihexadecyldimethyl ammonium chloride; amide derivatives of amino alcohols such as \(\beta-hydroxyethylstearyl-amide; amine salts of long chain fatty acids; quaternary ammonium bases derived from fatty amides of	35
40	di-substituted diamines such as oleyibenzylaminoethylene diethylamine hydrochloride; quaternary ammonium bases of the benzimidazolines such as methylheptadecyl benzimidazol hydrobromide; basic compounds of pyridinium and its derivatives such as cetylpyridinium chloride; sulfonium compounds such as octadecylsulfonium methyl sulfate; quaternary ammonium compounds of betaine such as betaine compounds of	40
45	diethylamino acetic acid and octadecylchloromethyl ether; the condensation products of stearic acid and diethylene triamine; polyethylene diamines; and polypropanol-polyethanol amines. The emulsifier is conventionally employed at a level of 1%—10% by weight of the siloxane, more preferably 0.5%—5% by weight.	45
50	The catalyst employed to polymerise the siloxane is preferably an alkaline catalyst such as an alkali metal hydroxide or a quaternary ammonium hydroxide of the formula (R°),N°OH ⁻ . In such ammonium hydroxides the R° groups can be hydrogen or alkyl radicals such as methyl, ethyl, propyl, butyl, isobutyl, decyl or octadecyl or aralkyl radicals such as benzyl or hydroxyalkyl radicals such as hydroxyethyl, hydroxypropyl	50
55	and hydroxybutyl.	55
J.	Most preferably the catalyst is a quaternary ammonium hydroxide having at least one radical of at least 12 carbon atoms in chain length, such a material also serving as an emulsification agent. Long chain length alkyl quaternary ammonium salts are also preferred as the emulsification agents, particularly di-long chain alkyl di-lower alkyl	
60	quaternaries, such as ditallowyl dimethyl ammonium chloride (DTDMAC), available commercially from Armour Chemical Company as Arquad 2HT (Arquad is a Registered Trade Mark) and imidazolinium derivatives such as methyl C _{1s} alkyl amidoethyl, C _{1s} alkyl imidazolinium methosulphate, available commercially from Ashland Chemical Company as Varisoft 475 (Varisoft is a Registered Trade Mark).	60

		<u> </u>
5	The level of catalyst usage is dependent on the catalyst type employed. Acid catalysts are conventionally used at high levels, e.g. at 15% or more by weight of the aqueous phase of the emulsion. Alkaline catalysts by contrast are used at lower levels, e.g. from 0.001% to 10%, preferably from 0.1% to 5% by weight of the siloxane monomer.	5
10	Emulsion polymerisation of dimethyl siloxane using DTDMAC as emulsifier In a typical preparation, dichloro dimethyl siloxane was first hydrolysed to form octamethyl cyclo tetra siloxane using the method of Patnode and Wilcock in JACS 68 1946 pp 358—363. 15 grs of this material were then added to a mixture of 131 grs of a 1% aqueous solution of ditallowlyl dimethyl ammonium choide and 3.75 grs of tetrabutyl ammonium hydroxide in the form of a 40% aqueous solution. The mixture	10
15	was stirred during addition of the ingredients by means of a Silverson (trade mark) laboratory emulsifier mixer and, after addition was complete, the reaction mixture was subjected to 15 minutes further agitation using an ultrasonic vibrator. After 18 hours at 80°C the emulsion was neutralised and the poly dimethyl siloxane oil was precipitated from the reaction mixture by addition of 500 mls of ethyl alcohol and was then dried with further alcohol before being heated at 75°C under a high vacuum	15
20	to remove all volatile materials. The viscosity of the silicone was determined to be 22,000 centistokes by measurement of its rate of flow under gravity between two marks on a calibrated tube. The time taken for a given quantity to flow along the tube was converted to viscosity using a calibration curve established with commercial silicones of known viscosity.	20
25	Using the above-described polymerisation technique, stable 10% emulsions of polydimethyl siloxane were achieved and equivalent results were obtained when the procedure was repeated using cetyl trimethyl ammonium bromide and the imidazoline derivative Varisoft 475 respectively as the emulsifier.	25
30	b) α-ω quaternised polysiloxanes The preparation of α,ω quaternised siloxane polymers can be conveniently carried out using the method disclosed in I.C.I. British Patent Specification No. 1,006,729. In this technique a polysiloxane end-stopped with alkyl halide groups in which the halogen atoms are separated from the nearest silicon atoms by at least three carbon atoms, is reacted with a tertiary amine to give an α,ω-quaternised siloxane polymer. In order to provide the polysiloxane starting material a solution polymerisation is normally	30
35	carried out to give a polymer of the appropriate molecular weight, and the polymerisation reaction is terminated by reaction with a whalo alkyl dimethyl silanol. As previously stated, the polysiloxane can be a poly di (C ₁ —C ₅ alkyl) or (C ₁ —C ₅ alkyl, aryl) siloxane, preferably a polydimethyl siloxane and the tertiary amine can be any alkyl, aryl or mixed alkyl and aryl material. Examples include trimethylamine, cetyl dimethyl-amine, phenyl dimethyl-amine.	35
40	Preparation of a and dipyridinium polydimethyl siloxane A typical preparation of this class of silicone polymers involved the polymerisation of 23.2 grs of octamethyl cyclotetra siloxane in the presence of 0.9 mls of concentrated sulphuric acid and 2.5 grs of 1,3-bis, 3-chloropropyl tetramethyl disiloxane. The	40
45	mixture was shaken in a sealed flask for 48 hours at room temperature, following which 5 mls of water were added and the flask shaken for a further hour. The resulting emulsion was split by addition of 50 mls of diethyl ether and the organic layer was then washed twice with 30 ml aliquots of distilled water, dried over sodium bicarbonate and magnesium sulphate and filtered. Evaporation of the filtrate to remove the ether left 23 much as all of interesting 100 m. NATA completion of the oil showed it to	45
50	left 23 grs of a clear oil of viscosity 100 cs. NMR examination of the oil showed it to correspond to a polymer having 36 siloxane units. 10 grs. of the a ₅ to-bis(3 chloropropyl) silicone prepared above were then refluxed in 10 mls pyridine for 36 hours at 120°C. Excess pyridine was distilled off under reduced pressure leaving a brown viscosit oil. This was then dissolved in tolurne,	50
55	washed with water and the toluene layer dried and evaporated to remove the toluene. NMR spectral analysis disclosed a level of proton activity corresponding to 70—80% of the theoretical uptake of pyridine. 10% aqueous emulsions of the silicone product were prepared by mechanical emulsification using an ethoxylated linear alcohol emulsifying agent (Dobanol* 45B4).	55
60	a C ₁₄ —C ₁₅ linear alcohol tetra ethoxylate supplied by Shell International Chemicals Limited) at a level of 20% by weight of the siloxane.	60

^{*} Dobanol is a Registered Trade Mark.

5	c) Amino functional linear polysiloxanes Amino functional linear polysiloxanes can be prepared by the general method disclosed in British Patent Specification No. 1,339,906 at page 3 lines 78—108, page 4 lines 1—65 and page 3 lines 3—14. In this method, a hydrosiloxane is reacted with an alkenyl group-containing tertiary amine in the presence of a platinum catalyst in accordance with the equation	5
	$(Me_3SiO)_2 (SiMe_2O)_x(SiMeHO)_y + yCH_2 = CHR'NR_2 \xrightarrow{H_2PtCl_0} (Me_3SiO)_2 (SiMe_2O)_x(OSi(Me)R''NR_2)_y$	
10	wherein x=10 to 100, y=1 to 20, R is a methyl, ethyl or phenyl group, R' is a direct linkage or a divalent organic group free of aliphatic unsaturation containing 1—16 carbon atoms and R'' is a divalent organic group free of aliphatic unsaturation containing 2—18 carbon atoms. The product of the above reaction can then be quaternised by further reaction with an alkyl halide or can be converted to the hydrochloride by acidification with hydrochloric acid.	10
15	Preparation of polydimethyl siloxane substituted with dimethylaminopropyl groups In a typical preparation 50 grs of dimethyl-methyl hydrogen siloxane copolymer containing approximately 76 dimethyl siloxane units and 6 hydromethyl siloxane units was dissolved in 50 mls toluene containing a trace of chlorplatinic acid. The mixture was stirred under nitrogen at 80°C, 5.18 grs of N,N-dimethyl allylamine in 10 mls of	15
20	toluene was added dropwise, holding the reaction temperature at 80—90°C, and the reaction mixture was stirred for a further 2 hours and then cooled. Sodium carbonate was added to neutralise any remaining acid and the mixture was filtered and rotary evaporated to remove solvents, leaving a pale yellow fluid of low viscosity. NMR analysis showed the formation of poly dimethyl silozane containing dimethylamino-	20
25	propyl groups at a level corresponding to a reaction completeness of 80%+, and a degree of substitution (d.s.) of 0.06. 20 grs of the reaction product was stirred in 100 mls of a 1:1 mixture of dichloromethane and isopropanol and 1.3 mls of concentrated HCI (11.21M) in 10 mls of the same solvent mixture was added slowly at room temperature. Following evaporation of	25
30	the solvent a pale coloured solid was left and NMR analysis showed this material as having a proton ratio close to the expected value for the hydrochloride derivative with no detectable level of the starting material. The siloxane polymer was then made up into a 10% aqueous emulsion using 20% based on the siloxane weight of a nonionic emulsification agent (Dobanol* 45E4, a linear C ₁₄ —C ₁₆ alcohol containing 4 moles of ethylene oxide supplied by Shell International Chemicals Limited).	30
35	A similar experimental technique to the above was employed to produce poly- dimethyl siloxanes having respectively approximately 40 siloxane units and a d.s. of 0.04 and 72 siloxane units with a d.s. of 0.015. The concentration of the aqueous dispersions (by which term is included solutions)	35
40	which constitute the compositions of the invention is not critical and is controlled by practical considerations. Accordingly, the dispersions should be concentrated enough not to be wasteful in transit costs, yet should be sufficiently aqueous to maintain the silicone component in emulsified form and fluid enough to disperse readily in a usage bath. Usually a content of from about 1% to 20%, especially about 3—10% by weight of components (a) and (b) together is convenient. Thus, the compositions can take the	40
45	form of an aqueous dispersion or solution of comparatively low viscosity (i.e. 200 centistokes or below) or of a paste, cream or gel. As stated earlier, the ratio of the siloxane portion of component (b) to the quaternary softening agent of component (a) should be in the ratio of from 5:1 to 1:100, preferably from 2:1 to 1:10 and most preferably from 1:1 to 1:3.	45
50	-	50
55	Optional Components The aqueous dispersions may contain other components, such as nonionic emulsifying aids, which are used at levels of the order of 1% by weight of the composition to aid dispersion of the usually poorly soluble cationic softeners. A wide range of nonionic emulsifiers can be used for this purpose such as those disclosed in German Patent Application OLS 2500111 published July 17th, 1975. It has also been	55

^{*} Dobanol is a Registered Trade Mark.

Colourists—Three dimensional durable press replicas for use with AATCC Test 124).

	-5-10,200		10				
	Number 8 graded perfect, Number 1 worst. A grade 5—7 was about that degree of freedom from wrinkling at which a housewito consider ironing unnecessary.	deemed to represent fe might be expected					
5	Ease of Ironing test This was judged by a panel of judges, employing a Scheffe gradings (panel score units -psu) and a "yardstick", i.e. least at 95% probability.	e analysis to provide difference significant	5				
	End Result Test A visual preference, graded as above in psu.						
10	Softness test A tactile preference, graded as above in psu.		10				
	Drying of Fabrics The "spin-dried" test pieces were dried by hanging in t drying) or in a tumbler dryer.	he laboratory (static					
15 20	Example I Comparison of silicones having different viscos The effect of silicone viscosity on the wrinkle grade, ease ironed-end-result grade of cotton tea towels treated with composi was examined for a range of polydimethyl siloxanes prepared by tion using a cationic emulsifier.	of-ironing grade and tions of the invention	15				
	Seven aqueous compositions [of which (A)—(D) were for comparison purposes, (E)—(G) being according to this invention] were made up, each containing 6% DTDMAC and 3% of a silicone emulsion polymerised in the presence of 1% DTDMAC on a silicone basis. The siloxane polymers varied in viscosity from 1000 to						
30	170,000 centistokes. The compositions were applied at 0.2% concesimulating a final rinse treatment in a conventional washing of fabrics were then air dried and graded by a panel of judges. In difference between fabrics of approximately 0.5 panel score ungrade is normally detectable by the housewife, while for ease of end result a panel score unit difference of between 0.75—1.1 generally necessary in order for a difference to be noticeable. The results are shown below, indexed to the grades obtain emulsified silicone of viscosity 170,000 centistokes.	ycle and the treated such panel testing a nits (psu) in wrinkle i ironing and ironing between fabrics is	25. 30				
35	Siloxane Wrinkle Ease of Irone Viscosity cs Grade Ironing end re (A) 170,000 0 0 0 (B) 100,000 +0.2 0 +0.5	sult	35				
40	(C) 40,000 +0.3 +0.1 -0.2 (D) 20,000 +0.4 +0.8 +1.2 (E) 8,000 +0.6 +0.7 +1.7 (F) 3,700 +0.6 +1.1 +2.1 (G) 1,000 +0.5 +0.8 +1.3		40				
45	It can be seen that performance for these parameters impro in viscosity from 170,000 cs, with the optimum appearing to lie 8,000 cs.	ves with a reduction in the range 3,000—	45				
50	Five compositions (A)—(E) in accordance with the invention tested for ease of ironing, wrinkle grade, ironed end result and supercially available fabric softening product containing 5.8% ammonium chloride as the standard. Positive figures denote an approduct and vice versa. An asterisk denotes a significant different of confidence.	oftness, using a com- ditallowyl dimethyl dyantage for the test	50				

1.	54	9.	1	R	n
4,	~~	,		U	v

11			1,549,180)				11
<u></u>	Formulation			Le	vels			
	Lommanon		(A)	(B)	(C)	(D)	(E)	
	N-tallowyl, N-methyl-N	'dimethyl-	• •					
	1,3-propane diamine							5
5	hydroacetate	r/ 1:tt	4	4	4			3
	N-tallowyl, N-methyl N	ramemyi-						
	1,3-propane diamine hydrochloride					4	4	
	Glyceryl monostearate		2	2	2	2.5	4.5	
10	Releasil 8 (a nonionic e	emulsified						10
	polydimethyl siloxane	of viscosity						
	1000 cs) available fro	m Dow	2	2	2	2	1.5 ²	
	Corning ("Releasil" is Unmodified White Dex	a mane mark)	L	L	3.0 ¹	_		
15	Cationic pyro dextrin	41414		2.9 ²				15
1.5	Ease of ironing		+0.78*	+ .82*	+1.2*	1.2*	+0.92*	
	Wrinkling		+0.32*	+0.04	-0.3*	+0.28	+0.02 +0.22*	
	Ironed end result		102	-0.10	-0.26*	+0.3* +0.56*	+1.24*	
	Softness		+0.3			T020	, ,,,,,,	
20	1. White dextrin	supplied by R	emy, Bel	gium.				20
	Reaction produ	ct of glycidyl	triethyl	ammoniur	n chloride	with whit	e dextrib	
	to give a d.s. o	of 0.85.	.c. 41		oilovona /	of viceocit	# 350 cs	
	3. Q2-1070—A n available from	Onionic emilis	inea bor	ушшешуг	SHOZAHE (OI ATSCOST	y 550 C	
25	It can be seen that	the composit	ions of th	ne inventio	n provide	an improv	rement in	25
2.5	eace of imping and W	rith one exce	ntion, an	improver	nent in w	mikie grac	ie or me	
	fabrics prior to ironing	and in irone	ed end re	sult. Soft	ness of the	treated in	abric was	
	also enhanced in those	instances whe	re this va	ariable wa	s measured	l.		
		E	KAMPLE	3 III			_	
30	A fabric treating c	omposition (A) in acco	rdance wit	th this inve	ention was	made up	30
	comprising, by weight,	ra C						
	3 part DTDM	Aqueous emu	Isian of 1	Polvdimetl	nvi siloxan	e of visco	sity 5000	
	centist	okes prepared	bv emu	lsion poly	merisation	in the pr	esence of	
35	10%	DTDMAC en	nulsifier (based on a	siloxane w	eight).		35
	1.50 " cation	ic pyrodextrin	compris	ing Britis	h Gum m	nodmed by	reaction f 0.07	
	with g	lycidyl trimet ic emulsifier	ryi ammo	misture o	onae w gr f Territol	trade ma	rk) 15S3	
	0.50 so nonion	5S9 secondary	-alcohol	ethoxylates	marketed	by Union	Carbide)	
40	0.35 " colour	perfume and	miscella	neous		•		40
	79.65 water.	_					f-b-i-	
	This composition	was compare	d with a	commerc	rally avail	able aquec	oult anti-	
	softener (B) containing wrinkling (i.e. appeara	5 6% DIDM	MC IOI	ease or m	oning, nor	rdv. on c	otton and	
45	-al-moston action bland	millouvesses (i	nen_coffo	n tea mwi	us and ter	гусоцоп ш	weis.	45
43	Each product was	applied in the	e final rii	nse of an	automatic (60°C. wasi	n cycle in	
	210H water, followed	by som drym	g and su	anc arying	z monors.			
	Test results are g	iven below. E	ase of iro	ning, iron	ed end res	zoro Anti	minkling	
50	reported in panel score	obsolute and	ee poorer	resunt ext	icoscu as	ZIV IIII	-MIIIIII	50
50	results are reported as				D:11	Test	owels	
		Cotton Pillos B A	wcases r 1sd	B A		B	A 1sd	
	Ease of Ironing	0.00 1.80	0.24	0.00 1.0			.20 0.94	
	Total of Houne	0.00 1.10	0.69	0.00 1.3	30 0.61		.40 0.66	<u> </u>
55	Ironed End Result	0.00 0.72	0.90	0.00 0.0			00 0.94	55
-		0.04 0.00	0.75	0.26 0.0 3.30 3.			0.00 1.08 2.75	
	Anti-wrinkling	1.85 1.90 1.25 1.35		3.30 3. 3.50 3.			i.40	
		1.22			Towels			
60	Cafeness			0.14 0.0				60
w	Softness			0.00 0.				

Feel/body was expressed in terms of preference of a 20-member panel.

				123200				12
			7	Test I		Tes	t II	
		_		. No	_		No	
	Comm31	R	A	preference	В	A	preference	
5	Cotton pillowcases	6	11	3	6	12	2	
J	Poly/cotton	7	8	4	9	9	2	5
	Tea Towels	. 6	12	2	1	16	3	
	Terry Towels	7	10	No preference 3 4 2 3	5	12	3	
10	It can be seen that consignificantly better Ease of advantage in Appearance between Result performance.	Ironii	ng and	Body/Feel be	nefits	as wel	l as an overall	10
15	aikyi chains c	position cation ono-ar or one ridazol	nic com nmoniu C ₁₈ —(inium t	pound selected m compounds 24 alkyl chain; extile softeners	from having	g either	on comprising: two C ₁₂ —C ₂₀	15
	R _B - N - (CH _Z R _B - N - (CH _Z						Э	
20	alkenyl group hydrogen, C, is ≪25; R, ii n is from 2 t	C; K ₁₆ Co defin o 6 ar	, K ₁₁ alkyl, ed as R nd X⊖∶	and C_{12} are in $(C_2H_4O)_pH$ or C_3 or as R_{10} , R_1 is an anion;	ndepe (C.H	ndently LO).H.	alkyl group or selected from wherein p+q is from 0 to 8,	20
25	(iv) compounds of	the	general	formula:				25
•	wherein R ₁₈ i	s hydr	ogen or	R _B R _B , (so	i grou	p. r is :	an integer from	
	(v) mixture there	of on	tuan 3	and X9 is an	mon;	and		
30	(b) an emulsion of a p siloxane in which th which may be subs a viscosity at 25°C	redom e alky tituted of at	inantly I group with c least 10	s may be partia ationic nitrogen O centistokes ar	lly or group id up	wholly ps, the to 8000	fluorinated and siloxane having centistokes:	30
35	the weight ratio of the silox the range from 5:1 to 1:10 2. A composition accord compounds has the formula	Ю.				_	_	35
		ъ	D D	D. NITH VIA				
	wherein X is an anion and	_	A-1 A-2	K² M⊕ X⊖				
40	(i) R and R ₁ each rep and R ₂ and R ₃ ar groups or R ₂ and	resent e inde R _o , t	ependen ogether	tly selected fro	nna Ca	Caal	kyl and benzyl	40
45	membered heteroc (ii) R represents a strai a C ₁ —C ₂ alkyl or 1 3. A composition accor softener has the general form	ght or benzyl rding	branch group	and R_2 and R_3 :	are as	hereina	bove defined.	45
	 -	ار ار ار	ե [⊕] — ၄ ₂ ,	4-n - C - R6	×Θ			

	wherein R_4 is hydrogen or a C_1 — C_4 alkyl group, R_5 is a C_1 — C_4 alkyl group, R_6 is a C_8 — C_{25} alkyl group, R_7 is a C_8 — C_{25} straight or branched chain alkyl group and X^{Θ}	
	is an anion.	
	4. A composition according to Claim 3 wherein the weight ratio of the siloxane	
5	content of component (b) to component (a) is from 2:1 to 1:10.	5
,	5. A composition according to any one of the preceding claims in which com-	
	ponent (b) comprises a linear poly di(C ₁ —C ₃ alkyl) siloxane prepared by emulsion	
	polymerisation using a cationic emulsifier.	
	6. A composition according to claim 5 wherein the cationic emulsifier is a fabric	
10		10
10	softening cationic surfactant.	10
	7. A composition according to any one of claims 1—4 wherein component (b)	
	is an α,ω-diquaternised linear di(C ₁ —C ₅ alkyl) siloxane polymer.	
	8. A composition according to claim 7 wherein the quaternising group comprises	
	an aromatic molecule.	
15	9. A composition according to any one of claims 1—4 wherein component (b)	15
	comprises a linear di(C ₁ —C ₃ alkyl) siloxane polymer in which from 0.001% to 0.1%	
	of the siloxane units contain an amino substituent group.	
	10. A composition according to claim 9 in which the degree of substitution (d.s.)	
	lies in the range from 0.01% to 0.075%.	
20	11. A composition according to any one of claims 1-4 wherein the component	20
	(b) is a polydimethylsiloxane.	
	12. A composition according to Claim 11 wherein the polydimethyl siloxane has a	
	viscosity at 25°C of 1,000—5,000 centistokes.	
	13. A composition according to any one of the preceding claims containing from	
25	1%-20% by weight of components (a) and (b).	25
	14. A fabric treating composition according to claim 13 containing from 3% to	
	10% by weight of components (a) and (b).	
	15. A fabric treating composition substantially as described with reference to any	
	one of the Examples I (E)—(G), Examples II (A)—(E) and Example III (A).	
30	16. A method of treating fabrics comprising the step of steeping them in an	30
	aqueous bath containing a composition as claimed in any one of claims 1-15 said	
	bath containing from 20-1000 parts per million (ppm) by weight of components (a).	
	and (b) together of which at least 10 ppm is component (b).	
	17. Fabrics when treated by the method according to claim 16.	
	_,,	

For the Applicants: CARPMAHLS & RANSFORD, Chartered Patent Agents, 43, Bloomsbury Square, London, W.C.1.

Printed for Her Majesty's Stationery Office, by the Courier Press, Leamington Spa, 1979
Published by The Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from
which copies may be obtained.